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UNITED STATES ENVIRONMENTAL PROTECTION AGENCY WASHINGTON, D.C. 20460

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OFFICE OF
PESTICIDES AND TOXIC SUBSTANCES

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MEMORANDUM

SUBJECT: EPA File Symbol 2935-UUN

Nu-Zone 10ME

FROM:

Mary L. Waller

Technical Support Section

Fungicide-Herbicide Branch

Registration Division (TS-767C)

TO:

Lois A. Rossi, PM 21

Fungicide-Herbicide Branch

Registration Division (TS-767C)

APPLICANT: Wilbur-Ellis Company

191 West Shaw Avenue, Suite 107

Fresno, CA 93704-2876

ACTIVE INGREDIENT:

Imazalil: (1-(2-(2,4-dichlorophenyl)-2-

(propenyloxy)-ethyl)-lH-imidazole 109

BACKGROUND:

The applicant has submitted an acute inhalation toxicity study and a dermal sensitization study as requested by the 10-31-86 TSS review of other acute toxicity data. The studies were conducted by Northview Pacific Laboratory, Inc. The MRID numbers are 401890-01 and -02. The method of support was not indicated.

RECOMMENDATION:

FHB/TSS finds the acute inhalation and dermal sensitization study acceptable to support registration of 2935-UNN. The signal word is CAUTION.

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LABELING:

Add the following sentences to the precautionary statements: "This product may cause an allergic skin reaction."

REVIEW:

(1) Acute Inhalation Toxicity Study: Northview Pacific Laboratory, Inc.; NVP Report number U7A009G; MRID number 401890-01; 3-4-87.

PROCEDURE:

Five male and five female Sprague-Dawley derived rats were exposed 4 hours in a 200 L chamber to a mean gravimetric concentration of 5.44 mg/L of test material. Animals were weighed prior to exposure and at 7 and 14 days. Animals were observed twice daily for the first week and once daily thereafter for 14 days. Animals were necropsied at study conclusion.

RESULTS:

No deaths occurred; therefore, it can be assumed that the LC50 is > 5.44 mg/L. Observations included atoxia, noisy respiration, and piloerection. Gross necropsy revealed slightly mottled lungs.

STUDY CLASSIFICATION:

Core Guideline Data

TOXICITY CATEGORY: Category IV - CAUTION

(2) <u>Dermal Sensitization Study</u>: Northview Pacific Laboratory; NVP Report number X7A069G; MRID number 401890-02; 2-20-87.

PROCEDURE:

A test group of ten Hartley guinea pigs and a positive control group of five Hartley guinea pigs received six induction treatments which were applied to a previously shaven site on each animal's shoulder and covered with occlusive wrap for six hours of exposure. Induction treatments were administered three times a week for two weeks as follows: the test group received 0.5 ml of undiluted test material and the positive control group received 0.5 ml of a 0.1% solution of l-chloro-2,4-dinitrobenzene. Animals were placed in restrainers during exposure. Animals' shoulders were cleaned after each exposure and redepilated as needed. Two weeks

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after the last induction treatment, animals were challenged for 24 hours at virgin sites. Skin irritation was scored at 24 and 48 hours.

RESULTS:

All test animals exhibited moderate erythema to erythema and cracking of the skin during induction. The positive control animals exhibited slight confluent erythema to edema and cracking of the skin during induction. At challenge, both the test animals and positive control group exhibited moderate patchy erythema to erythema with edema and cracking of the skin.

STUDY CLASSIFICATION: Core Guideline Data

TOXICITY CATEGORY: Sensitizer

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	Identity of product impurities
	Description of the product manufacturing process
	Description of product quality control procedures
	Identity of the source of product ingredients
	Sales or other commercial/financial information
<u>x</u>	A draft product label
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